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A pharmaceutical composition comprising a suitable pharmaceutical vehicle and an element chosen from the group consisting of a nucleotide sequence encoding a peptide of the CNECUT family, a vector comprising this nucleotide sequence, the polypeptide sequence encoded by this nucleotide sequence and/or a cell line transformed with said vector and expressing the peptide of the ONECUT family.

- 10 2. The pharmaceutical composition as claimed in claim 1, characterized in that the peptide of the family is HNF-6.
  - The pharmaceutical composition as claimed in claim 1, characterized in that the peptide of the family is OC-2.
    - 4. The cellular pharmaceutical composition as claimed in claim 1, characterized in that the peptide of the family 18 OC-3.
  - 5. The pharmaceutical composition as claimed in any one of the preceding claims, characterized in that said nucleotide and polypeptide sequences are human nucleotide and polypeptide sequences.
  - The pharmaceutical composition as claimed in any one of the preceding claims, characterized in that the vector is chosen from the group consisting of plasmids, viruses, phagemids, lipid vesicles, in particular cationic vesicles, liposomes or a mixture of
  - these.
    7. The use of the pharmaceutical composition as
    7. The use of the praceding claims, for
    claimed in any one of the praceding claims, for
    preparing a medicinal product intended for the
    prevention and/or for the treatment of type 1 or type 2
    diabetes or of disorders linked to diabetes, for the
    prevention and/or for the treatment of cancer, in
    particular of melanoma, and for the prevention and for
    the treatment of Waardenburg syndrome.
    - A method of therapeutic treatment of a patient, preferably of a human patient, likely to develop or

suffering from diabetes, from a cancer, in particular from a melanoma, or from Waardenburg syndrome, characterized in that the pharmaceutical composition as claimed in any one of claims 1 to 4 is administered ex vivo by isolating a body fluid or one cr more cells from the patient, treating said cells or the cells present in this body fluid with the pharmaceutical composition of the invention or with the vector included in this pharmaceutical composition, and reinjecting into said patient the transformed cells.

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## CLAIMS

- A pharmaceutical composition comprising An acceptable pharmaceutical vehicle and an element chosen from the group consisting of a nucleotide sequence 5 enooding a protein of the ONECUT family characterized by the presence of a single CUT domain and the presence of an F48M50 dyad in the homeo domain, a vector comprising this nucleotide sequence, the protein sequence encoded by this nucleotide sequence and/or a cell line transformed with said vector and expressing 10 said protein of the CNECUT family.
  - The pharmaceutical composition as claimed in claim 1, characterized in that the protein of the ONECUT family is HNF-6 in its two isoforms.
- The pharmaceutical composition as claimed in claim 1, characterized in that the protein of the 15 CNECUT family is OC-2, the amino acid sequence of which is SEQ ID No. 2.
- The cellular pharmaceutical composition as claimed in claim 1, characterized in that the protein of the ONDCUT family is OC-3, the amino acid sequence 2.0 of which is SEQ ID No 3.
  - The pharmaceutical composition as claimed in any one of the preceding claims, characterized in that said nucleotide and polyreptide sequences are human nucleatide and polypeptide sequences.
  - The pharmaceutical composition as claimed in any one of the preceding claims, characterized in that the vector is chosen from the group consisting of plasmids, viruses, phagemids, lipid vesicles, in particular cationic vesicles, liposomes or a mixture of these.
  - The use of the pharmaceutical composition as claimed in any one of the preceding claims, for 35 preparing a medicinal product intended for the prevention and/or for the treatment of type 1 or type 2 diabetes or of disorders linked to diabetes, for the

prevention and/or for the treatment of cancer, in particular of melanoma, and for the prevention and for the treatment of Waardenburg syndrome.

8. A method of therapeutic treatment of a patient,
5 preferably of a human patient, likely to develop or
suffering from diabetes, from a cancer, in particular
from a melanoma, or from Waardenburg syndrome,
characterized in that the pharmaceutical composition as
claimed in any one of claims 1 to 4 is administered ex
10 vivo by isolating a body fluid or one or more cells
from the patient, treating said cells or the cells
present in this body fluid with the pharmaceutical
composition of the invention or with the vector
included in this pharmaceutical composition, and
15 reinjecting into said patient the transformed cells.

AMENDED PAGE